

(12) United States Patent

Malanowska-Stega et al.

(10) Patent No.:

US 9,351,712 B1

(45) **Date of Patent:**

May 31, 2016

(54) SIMULTANEOUS MULTIPLE METHOD **OUT-PATIENT UTERUS BIOPSY DEVICE** AND METHOD

- (71) Applicants: Zanetta Malanowska-Stega, Bronx, NY (US); Giuseppe DelPriore, Indianapolis, IN (US)
- (72) Inventors: Zanetta Malanowska-Stega, Bronx, NY (US); Giuseppe DelPriore, Indianapolis,
- (*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.
- (21) Appl. No.: 14/582,484
- (22) Filed: Dec. 24, 2014

Related U.S. Application Data

- (63) Continuation of application No. 14/010,862, filed on Aug. 27, 2013, now Pat. No. 8,920,336, which is a continuation of application No. 13/733,934, filed on Jan. 4, 2013, now Pat. No. 8,517,956, which is a continuation of application No. 12/639,600, filed on Dec. 16, 2009, now Pat. No. 8,348,856.
- (60) Provisional application No. 61/138,055, filed on Dec. 16, 2008.
- (51) Int. Cl. A61B 5/00 (2006.01)A61B 10/02 (2006.01)A61M 25/09 (2006.01)
- (52) U.S. Cl.

CPC A61B 10/0283 (2013.01); A61B 10/0291 (2013.01); A61M 25/09 (2013.01); A61B 2010/0208 (2013.01); A61B 2010/0216 (2013.01); *A61M 2025/09191* (2013.01)

(58) Field of Classification Search

CPC A61B 10/02; A61B 10/0283; A61B 10/0291; A61B 2010/0216 See application file for complete search history.

(56)References Cited

U.S. PATENT DOCUMENTS

4,227,537	A *	10/1980	Suciu A61B 10/0291 600/569
4,324,262	A *	4/1982	Hall A61B 10/02
4,361,948	A *	12/1982	Omata A61B 10/02
4,485,824	A *	12/1984	29/517 Koll A61B 10/04
4,662,381	Α	5/1987	435/304.1 Inaba
6,036,658	A	3/2000	Leet et al.
8,348,856	B1*	1/2013	Malanowska-
			Stega A61B 10/0291 600/565
8.517.956	B1*	8/2013	Malanowska-
-,,			Stega A61B 10/0291 600/565
8,920,336	B1 *	12/2014	Malanowska-
			Stega A61B 10/0291 600/565
2002/0161313	A1	10/2002	Sak
2004/0116827		6/2004	Tiberio
2008/0188769		8/2008	Lu

^{*} cited by examiner

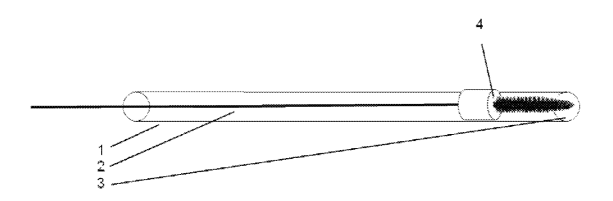
Primary Examiner — Max Hindenburg

(74) Attorney, Agent, or Firm — Steven M. Hoffberg, Esq.; Ostrolenk Faber LLP

ABSTRACT

A biopsy device, comprising a flexible coaxial structure, comprising an obturator within a sheath, the obturator being adapted to be displaced with respect to the sheath along the coaxial axis by a force applied at a proximal end; a disruptor, at a distal end of the obturator, adapted to disrupt a tissue surface to free cells therefrom, having a first position covered within the sheath and a second position freely extending beyond the sheath; an element, having a fixed position on the obturator, and a close clearance with respect to an inner surface of the sheath, such that a retraction of the obturator within the sheath causes an associated change in pressure in a space between the obturator distal to the element and the sheath, to thereby draw fluid proximate to an end of the sheath within the sheath.

20 Claims, 5 Drawing Sheets



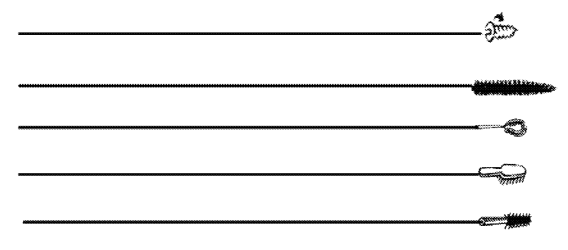
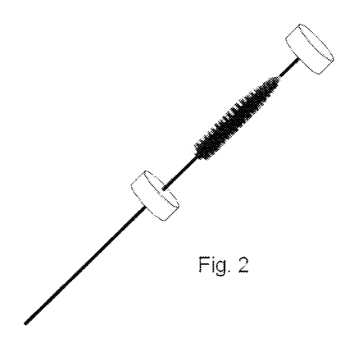
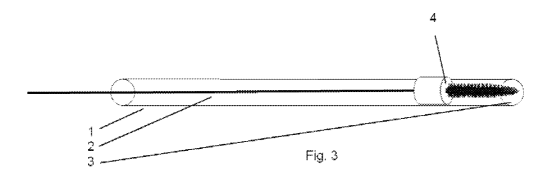


Fig. 1





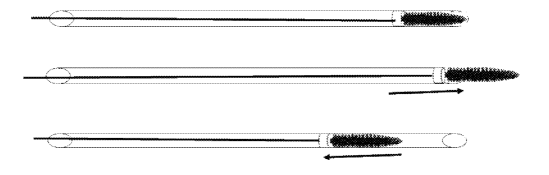


Fig. 4

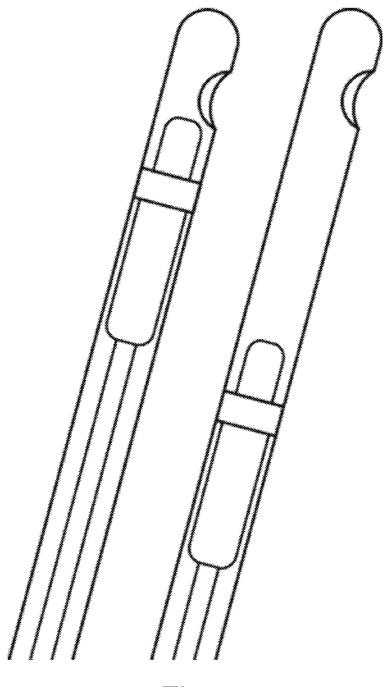


Fig. 5

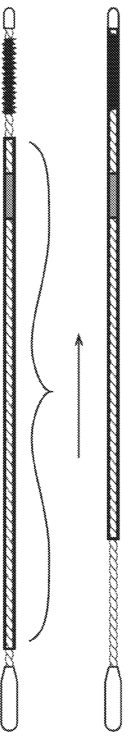


Fig. 6

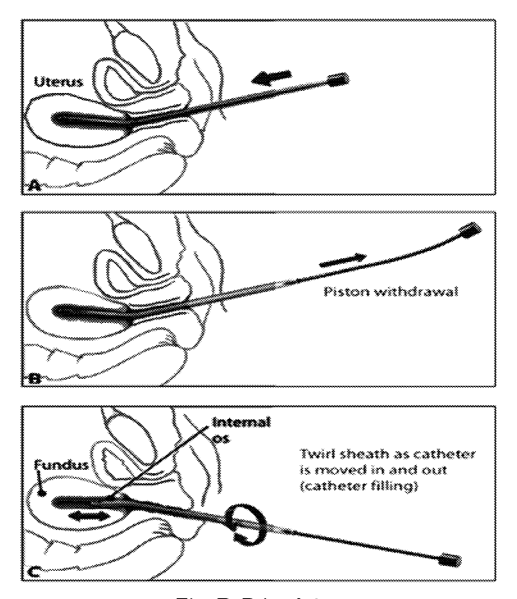


Fig. 7 Prior Art

SIMULTANEOUS MULTIPLE METHOD OUT-PATIENT UTERUS BIOPSY DEVICE AND METHOD

CROSS REFERENCE TO RELATED APPLICATIONS

The present application is a Continuation of U.S. patent application Ser. No. 14/010,862, filed Aug. 27, 2013, now U.S. Pat. No. 8,920,336, issued Dec. 30, 2014, which is a Continuation of U.S. patent application Ser. No. 13/733,934, filed Jan. 4, 2013, now U.S. Pat. No. 8,517,956, issued Aug. 27, 2013, which is Continuation of U.S. patent application Ser. No. 12/639,600, filed Dec. 16, 2009, now U.S. Pat. No. 8,348,856, issued Jan. 8, 2013, which is a non-provisional of U.S. Provisional Patent Application 61/138,055, filed Dec. 16, 2008, both of which are incorporated herein by reference in their entirety.

FIELD OF THE INVENTION

The present invention provides a system and method for performing a biopsy of the uterus. More particularly, it is a mechanical device that performs a mechanical disruption of the uterus, especially the endometrium, and simultaneously 25 aspirates the residue of the disruption.

BACKGROUND OF THE INVENTION

Current biopsy devices and methods do not consistently 30 provide an adequate sample to diagnose uterine abnormalities. For instance, if a cancer exists on the inside of the uterus, i.e. the endometrium, current devices and methods may miss the cancer and preclude an accurate diagnosis. To ensure an adequate and accurate specimen currently often requires a 35 painful and dangerous series of procedures.

For example, the Cook Medical Tao Brush™ I.U.M.C. Endometrial Sampler, and the Pipelle endometrial biopsy device (See, Sierecki A R, Gudipudi D K, Montemarano N, Del Priore G., "Comparison of endometrial aspiration biopsy 40 techniques: specimen adequacy." *J Reprod Med.* 53(10):760-4, 2008 October), expressly incorporated herein by reference.

The currently available options fall into two categories. The first are devices and methods that can be used in the office. These are usually thin devices 2-5 mm diameter that 45 rely on either a weak suction or a disruption of the uterus. These are relative safe options but are not considered sufficiently accurate. The other category consists of larger devices that require more analgesics and even an operating room. This category includes the traditional dilation and curettage, i.e. 50 D&C, and hysteroscopy. This category is more dangerous and painful than the first category. Neither category consistently gives an adequate, accurate and safely obtained sample and diagnosis in every case.

Endometrial sampling has become an integral part of the 55 workup of pre-menopausal and postmenopausal women who are having abnormal uterine bleeding. Approximately over one million endometrial sampling procedures are done in the United States annually. Of the women born today, 2.45% (1 in 41) will be diagnosed with cancer of the uterus sometime 60 during their lifetime. Therefore, theoretically 1 in 41 women will need an endometrial biopsy sometime in their lifetime.

Despite the large number of endometrial biopsies performed, limited information is available on the best technique to use with an aspiration type endometrial biopsy device (e.g. Pipelle, Cooper Surgical, Inc. Trumbull, Calif.). A slight difference in sensitivity and specificity of any screening proce-

2

dure will have a great impact on the general population in identifying cancer especially for a test performed so frequently.

Specimen adequacy is the important factor for a procedure to be accurate in detecting abnormal pathology, especially a potential malignancy, precancerous or early cancer lesion. Different endometrial biopsy techniques are used with the aspiration device; e.g. "cork-screw", "D&C", or a combination of both with little objective data on what technique is best.

Most tests have an inherent sensitivity and specificity. However alteration in technique can be an important part of the test performance. We noticed that some members of our faculty group practice had different techniques based on intu15 ition, e.g. povidone-iodine cleansing before endometrial biopsy. A literature reviewed and survey of the participating clinicians could not come up with a published evaluation of the different aspiration biopsy techniques used. We felt that the potential influence of these different techniques on the test characteristics of a test done more than one million times each year was important enough to begin an investigation of this area using a convenient sample and a retrospective chart review.

Based upon our analysis, the combined technique is significantly better than using either technique alone. Due to the large number of endometrial biopsies performed each year, even a small difference in test characteristics can have significant clinical ramifications.

Endometrial biopsy in the office setting, which consists of endometrial sampling using an aspiration biopsy device, has been shown to have sensitivity equivalent to that of dilatation & curettage (D&C). Its advantages include lower cost, it does not require anesthesia and it can be carried out in an office setting. Analyses have shown that a D&C is more costly and no more accurate in diagnosing endometrial abnormalities than the aspiration biopsy device. Aspiration biopsy device has been shown in repeated controlled clinical trials to be as effective as a D&C in diagnosing malignancy.

Although the aspiration biopsy device can detect cancer accurately, they may not able to detect tumors localized to a polyp or a small area of endometrium. Because of this variable and others, the sensitivity of an aspiration biopsy device in detecting uterine cancer varies in studies from 67% to 100% with an overall specificity of 100%. Huang et al. study found that an aspiration biopsy device had a sensitivity of 99.2% in pinpointing high-grade cancer and a sensitivity of 93% in detecting low-grade malignancies; the sensitivities as defined for D&C were 100% and 97% respectively. Low cost and accuracy dictate that an office aspiration biopsy device should be used as the frontline test for endometrial sampling and the detection of endometrial cancer.

The National Cancer Institute (NCI, 2008) has stated that there is insufficient evidence to establish whether a decrease in mortality from endometrial cancer occurs with screening by endometrial sampling. The NCI notes that based on solid evidence, endometrial biopsy (sampling) may result in discomfort, bleeding, infection, and in rare cases uterine perforation. In addition, risks associated with false-positive test results include anxiety and additional diagnostic testing and surgery. Furthermore, endometrial cancers may be missed on endometrial sampling.

Endometrial sampling by means of biopsy for histological examination in the diagnostic evaluation of abnormal uterine bleeding in women suspected of having endometrial hyperplasia or endometrial carcinoma is a minimally invasive alternative for dilatation and curettage (D&C) or hysteroscopy. The Pipelle endometrial sampling device is the most popular

method for sampling the endometrial lining (Guido, 2008). Various types of brushes have also been used for endometrial sampling. Although the brush appears to be as effective or better than other blind methods of endometrial sampling. these devices have been evaluated in only a few studies with small numbers of subjects (Critchley, et al., 2004; Yang, et al., 2002; Del Priore et al., 2001; Yang & Wan, 2000; Maksem, et al., 2000). In one of the larger comparative trials, 101 women (aged 35 to 86 years) with clinical indications for endometrial biopsy underwent a brush biopsy (Tao Brush, Cook OB-GYN, Bloomington, Ind.) and a Pipelle biopsy (Cooper Surgical, Shelton, Conn.) during one office visit. Twenty-two had cancer or atypia, the others had benign diagnoses. When correlated with the final diagnosis, sensitivity for the Tao Brush and Pipelle were 95.5% and 86%, respectively, and specificity was 100% for both (Del Priore, et al., 2001).

According to the company's website (Gynecor[™], Glen Allen, Va.), the TruTest[™] for total uterine testing is the first test that is able to detect endometrial and cervical cancer, 20 HPV, *chlamydia* and gonorrhea from the same specimen. Using the Tao Brush, a sampling of the uterine lining is taken and the brush is sent to Gynecor[™] for both histology and cytology examination. The testing kit provided by Gynecor has one Tao Brush (used for the collection of endometrial 25 tissue) and two cytobrushes (one is used to clean mucus and debris from the cervix and the second is used for enhanced cell and tissue collection from the squamo-columnar junction of the uterine cervix). The Tao Brush is an FDA, Class II device.

The use of histology for endometrial examination depends on having enough tissue to yield an accurate test result. However, a tissue specimen is sometimes hard to collect, especially in post-menopausal women. Gynecor fixative can be used for both histology and cytology. According to 35 Gynecore's website, "Cytologies are very important because they add about 20% more information than is obtained with just the histology. Using this method, Gynecor has been able to diagnose ovarian carcinoma in transit, endometrial intraepithelial neoplasia and endometrial intraepithelial carcinoma."

In a feasibility study, Maksem, et al. (1997) compared the cytologic diagnosis to the histologic diagnosis of endometrium collected from 100 hysterectomy specimens using the Tao Brush and the CytoRich fixative system. Inter- 45 pretative algorithms that translate histopathologic to cytopathologic diagnoses were used. The authors reported that cytology separated benign endometrium, low-grade (nonatypical) hyperplasia, high-grade (atypical) hyperplasia/ FIGO Grade I adenocarcinoma, and higher-grade carcinomas 50 from one another. Endometrial atrophy was diagnosed in 3 patients whose histology showed clinically asymptomatic, benign fibrous endometrial polyps. A low volume of abnormal cell aggregates interpreted as endometrial intraepithelial carcinoma was detected in 1 patient whose initial histology 55 was reported as simple hyperplasia, but whose histology on review after p53 staining revealed intraepithelial surface cancer. In the remaining 96 cases, the cytologic diagnosis consistently represented the histologic diagnosis of the hysterectomy specimen. On a case-by-case basis, any one cytology 60 slide accurately represented the diagnosis of the other cytology slides. The authors concluded that endometrial brushing with suspension fixation is (i) uniform, (ii) three-dimensional structures among cell aggregates are preserved, which allows pattern-based histologic diagnostic criteria to be applied to 65 cytologic samples, and (iii) only a limited number of slides need to be examined.

4

Van den Bosch, et al. (1998) evaluated the value of cervical cytology in menopausal women at high risk for endometrial disease in 128 consecutive menopausal women presenting with uterine bleeding (n=116) or in whom endometrial cells were found on a previous cervical cytology smear (n=12). An endo- and ecto-cervical smear was taken before hysteroscopy with curettage and the results of the cervical cytology were compared with the endometrial histology. Endometrial carcinoma was diagnosed by endometrial sampling in 6 women. In 2 of these cases cervical smears did not contain endometrial cells. The presence of endometrial cells on ecto-cervical cytology showed a sensitivity of 67% and a specificity of 78% for endometrial carcinoma versus 80% and 76%, respectively, for endo-cervical cytology. The positive predictive value for endometrial malignancy of the presence of endometrial cells on cervical cytology ranged between 13% and 17%. The presence of atypical endometrial cells on cervical smear was associated with endometrial malignancy in almost half the cases. The authors concluded that cervical cytology is of limited value in the diagnosis and the management of postmenopausal endometrial disease.

Dijkhuizen, et al. (2000) performed a meta-analysis to assess the accuracy of endometrial sampling devices in the detection of endometrial carcinoma and atypical hyperplasia. The authors searched the literature for studies published between 1966 and 1999 that compared the results of endometrial sampling with findings at D&C, hysteroscopy, and/or hysterectomy. They found 39 studies that included 7,914 women. For each study, the number of patients in which endometrial sampling failed as well as the sensitivity and specificity for the detection of endometrial carcinoma and atypical hyperplasia was calculated. The detection rate for endometrial carcinoma was higher in post-menopausal women compared with pre-menopausal women. In both postmenopausal and pre-menopausal women, the Pipelle was the best device, with detection rates of 99.6% and 91%, respectively. For the detection of atypical hyperplasia, there was only one study that reported explicitly on post-menopausal women, thereby hampering the possibility of subgroup analysis. Again, the Pipelle was the most sensitive technique with a sensitivity of 81%. The specificity of all devices was greater than 98%. The authors concluded that endometrial biopsy with the Pipelle is superior to other endometrial techniques in the detection of endometrial carcinoma and atypical hyperplasia. The accuracy of the Pipelle is higher in post-menopausal women compared with pre-menopausal women.

Maksem (2000) reported performance characteristics of the ability of the Tao Brush in recognizing histological patterns in cytology preparations of endometrial brushings (n=113). Correlative tissue examinations comprising Pipelle (Prodimed, Neuilly-en-Thelle, France) biopsy, hysteroscopy and biopsy, D&C, and hysterectomy were available at for 59 cases. In 42 cases, cytology diagnoses could be compared to histology diagnoses. Twenty-five of 63 normal brushings were followed up; 14 were normal. Eleven Pipelle biopsies of cytologically atrophic endometrium were quantitatively limited and insufficient for diagnosis. Thirty-seven cases were abnormal, and 15 of these showed nuclear anaplasia. Twentyeight of the abnormal cases were followed up. All correlative tissue examinations confirmed an abnormality. All 15 cases with nuclear anaplasia showed significant histopathology comprising atypical endometrial hyperplasia, endometrial intraepithelial neoplasia (EIN), endometrial intraepithelial carcinoma (EIC), and invasive adenocarcinoma. There were 13 inadequate endometrial brushings. Three cases had insufficient cellular material. The remaining 10 cases were cellular but were mainly cervical/endo-cervical samples. Two of the

cellular cases resulted from clinicians failing to replace the protective sheath over the brush bristles before removing the Tao Brush from the endometrial cavity. The remaining 11 cases resulted from inaccessibility of the uterine cavity due to a tight or stenotic cervix. The author concluded that (i) the Tao 5 brush is a reliable uterine sampling device for outpatient assessment of the endometrium of women with patent cervices, (ii) endometrial cytology accurately represents atrophic endometrium, (iii) it is an effective case-finding tool for EIN and EIC, and (iv) women with tight or stenotic cervices are 10 poor candidates for endometrial brushing, and may experience pain if the procedure is attempted.

In a case series on the use of the Tao Brush for endometrial biopsy, Wu, et al. (2003) reported that the sensitivity and specificity in identifying endometrial cancer was 100% and 15 96%; however, diagnosis relied mainly on histologic evaluation of hematoxylin and eosin-stained tissue sections and assessment of specimen adequacy was important when interpreting Tao Brush biopsies.

In an unblinded randomized trial, Critchley, et al. (2004) 20 compared 3 outpatient methods of endometrial evaluation in terms of performance, patient acceptability and cost-effectiveness. Women referred for investigation and management of abnormal bleeding between January 1999 and May 2001 were evaluated using blind biopsy alone, hysteroscopy with 25 biopsy, ultrasound evaluation including transvaginal ultrasound, and, in the low-risk group, the option of no investigation. Within this design, two devices for obtaining endometrial biopsy were compared, the Pipelle sampler and the Tao Brush. Minor adverse events (e.g., shock, patient distress) did 30 not occur for ultrasound, but occurred in 16% and 10% of women for hysteroscopy and biopsy procedures respectively. Pipelle biopsy provided an acceptable endometrial sample for 79% of moderate-risk women, but only 43% of high-risk women. The Tao Brush gave similar performance in moder- 35 ate-risk women (77%), but was more successful than the Pipelle sampler in post-menopausal (high-risk) women (72%).

To determine the performance characteristics of endometrial cytology for the detection of malignancy and atypical 40 hyperplasia using liquid-based cytology specimens collected with the Tao Brush sampler, Kipp, et al. (2008) obtained brushings of the endometrial cavity from 139 hysterectomy specimens before routine histopathologic evaluation. Cytology specimens were fixed in PreservCyt and processed using 45 ThinPrep technology. Cytology diagnoses were classified as non-diagnostic, negative, atypical, or positive for malignancy. Histopathologic findings were used as the gold standard for determining the performance characteristics of cytology. Histopathologic results from the 139 patients included 50 81 (58%) endometrial cancers, 7 (5%) complex hyperplasias with atypia, 2 (1%) complex hyperplasias without atypia, and 49 (35%) patients with benign histology. The number of specimens diagnosed cytologically as positive, atypical, negative, or non-diagnostic was 60 (43%), 40 (29%), 37 55 (27%), and 2 (1%), respectively. The overall sensitivity and specificity of cytology for detecting endometrial cancer and atypical hyperplasia were 95% and 66% when atypical cytology specimens were considered positive. The authors concluded that direct endometrial sampling by liquid-based 60 endometrial cytology collected with the Tao Brush sampler produces specimens that contain cellular material that may be identified as endometrial cancer or atypical hyperplasia; however, both atypical and positive cytology diagnoses are indicators for triage to more specific methods of diagnosis.

Williams, et al. (2008) evaluated factors affecting the adequacy of pipelle and Tao Brush endometrial sampling.

6

Women referred to an outpatient clinical for assessment of abnormal vaginal bleeding (n=200) were assigned to one of two risk groups: "high risk" for post-menopausal women and "moderate risk" for pre-menopausal women aged 40 years or older or with other risk factors. Women in each risk group had both Tao Brush and Pipelle biopsy and were then randomized to have either hysteroscopy and/or transvaginal ultrasound. Nulliparity was associated with failed insertion for both the Tao Brush and Pipelle (p<0.001). Among post-menopausal women, inadequate samples were associated with the Pipelle (p<0.001). Among pre-menopausal women with nulliparity, both the Tao Brush and Pipelle were associated with inadequate samples (p<0.001). A significantly greater proportion of women preferred the Tao Brush to the Pipelle.

Outpatient endometrial biopsy has a high overall accuracy in diagnosing endometrial cancer when an adequate specimen is obtained. A positive test result is more accurate for ruling in disease than a negative test result is for ruling it out. Therefore, in cases of abnormal uterine bleeding where symptoms persist despite negative biopsy, further evaluation will be warranted (Clark, et al., 2002). If the woman is postmenopausal and bleeding has not been persistent; a thin endometrial stripe in this setting is most consistent with atrophy and does not require further invasive studies. A thick endometrial stripe, persistent bleeding, or bleeding in a postor peri-menopausal woman should be followed by additional endometrial sampling, such as hysteroscopy with curettage. In asymptomatic post-menopausal women, the decision to biopsy is also based upon a variety of factors, including cervical cytology showing endometrial cells or glandular abnormality, and risk factors for endometrial cancer, such as unopposed estrogen and tamoxifen use. In a completely asymptomatic post-menopausal woman with no risk factors and an endometrial stripe less than 5 mm, there is no need for biopsy. However, even one drop of blood in a post-menopausal woman not on hormone therapy constitutes a symptom and is an indication for biopsy.

Current evidence-based guidelines from leading medical professional organizations include no recommendation for endometrial or cervical cytology performed in conjunction with endometrial histology in the diagnostic evaluation of abnormal uterine bleeding in women suspected of having endometrial hyperplasia or endometrial carcinoma. There is insufficient evidence to support this approach.

The problems presented in existing tissue sampling devices are solved by the device and method of the present invention.

SUMMARY OF THE INVENTION

A preferred embodiment of the present invention provides a narrow cylindrical tube with a guidewire and biopsy sampling device at the end, similar to a Cook Medical (Bloomington, Ind.) Tao BrushTM I.U.M.C. Endometrial Sampler, modified such that surrounding the guidewire, proximal to the biopsy sampling device at the end of the guidewire, and disposed within the sheath, is a piston-like structure which, when the wire is withdrawn through the sheath, draws a vacuum and sucks fluid surrounding the guidewire into the sheath. A vacuum biopsy sampling device, such as the known Pipelle endometrial suction curette produces a vacuum and draws it into the sheath by a similar principle, but lacks the brush or other biopsy sampling device at its distal end.

The preferred device is a 1-3 mm diameter by 30-40 cm long coaxial "straw" that can easily pass into the uterus endometrial cavity with little or no discomfort. It is malleable but rigid enough to apply sufficient force to pass through the

cervix. In the center of the outer sheath, which is an impermeable tube, a thinner inner insert can be extended beyond the end of the tube into the uterus. This inner obturator disrupts the uterus to loosen and collect a biopsy sample of the uterus.

The tissue sampling device includes a spirally twisted flex- 5 ible wire with opposed proximal and distal ends. Also included is a plastic tube covering a significant portion of the wire to provide additional rigidity without making the overall brush stiff. Along the distal end portion of the wire is a brush that includes bristles that were used for collecting a tissue 10 sample. The bristles are fixed within the spirally twisted wire near the distal end and are tapered from smaller to larger towards the distal end of the wire. Tapering of the bristles from the distal end of the device allows for more global tissue collection of the endometrium because of the shape of the endometrial cavity. An atraumatic bulb is located on the extreme distal end of the twisted wire.

Simultaneously with withdrawal of the inner obturator back into the narrow cylindrical tube, the device creates a weak suction to collect the disrupted sample into the outer 20 tube. The entire apparatus is then withdrawn from the uterus and the sample is collected by reversing the process outside the body.

In accordance with another embodiment, the plastic tube and twisted wire are contained within a sheath of shorter 25 need analgesics or an operating room. It is a safer, faster and length than the twisted wire, such that the sheath can be moved along the plastic tube to the atraumatic bulb on the distal end of the twisted wire, thereby covering the brush during insertion and removal after tissue collection. Before insertion, the sheath can be moved into position over the distal 30 end of the twisted wire to protect the brush during insertion. Having the brush covered during insertion also increases comfort for the patient and protects the brush from collecting tissue from unintended areas. The sheath is moved back toward the proximal end of the twisted wire after the device 35 has been inserted to the proper collection depth, exposing the brush and allowing for collection of a tissue sample. The sheath may be moved to completely uncover the brush or may be moved in gradients to uncover portions of the brush. This allows the practitioner to adjust the effective collection area 40 of the brush based on the anatomy of the patient. The plastic tube covering the wire is scored in centimeter gradations along the plastic tube with markings indicating the exact length of the brush inserted into the uterus, starting from the distal tip of the brush to the proximal end of the plastic tube. 45 Dijkhuizen F P, Mol B W, Brolmann H A, Heintz A P. The This allows the clinician to know how deeply the brush is inserted into the uterus. The sheath is approximately the same length as the plastic tube and in position to cover the brush bristles prior to insertion. The sheath may be formed of a clear material such that the gradations on the plastic tube may be 50 viewed through the sheath. The ability to measure insertion depth increases the certainty that the tissue sample collected is from the correct area, after a tissue sample is collected from the proper area, while the tissue sampling device remains inserted, the sheath can be moved back along the distal end of 55 the twisted wire to cover the brush bristles before removing the brush. This allows for the tissue sample to be protected on the brush within the sheath during removal.

Additionally, the gradations along the flexible tube allow the practitioner to measure the length of bristles exposed. As 60 the practitioner pulls the sheath from its insertion position towards the handle, the further the sheath is pulled the more bristles are exposed. The gradations (ruler) provide a visual confirmation of this measurement and allow the practitioner to be precise in exposing only a certain length of the brush bristles. This measurement allows the practitioner to have better control of where the tissue is sampled and allows the

8

practitioner to adjust the length of brush based on patient specific parameters; such as uterine size measured during previous tests or inferred based on patient history. Control of brush exposure increases sampling precision and patient

By combining two or more methods into one device, the invention eliminates pain, discomfort, and inconvenience, e.g., a second procedure to obtain an adequate and accurate specimen. The multiple methods of specimen collection, e.g., disruption by physical means, and suction, used together, allows a gentler application of the individual methods, e.g. a gentle disruption and gentle suction applied simultaneously can replace a vigorous disruption, e.g. D&C, and a powerful suction. The combination of multiple gentler methods in one device is safer and more effective than any method alone. On the other hand, to the extent either of the multiple methods is acceptably used at the levels used when employed individually, the apparatus and method according to the present invention has the capacity to produce improved results. The simultaneous combination of multiple methods provides a consistently more satisfactory specimen and more accurate diagnosis.

The invention can be used in a doctor's office and does not less expensive improvement over current methods.

REFERENCES (INCORPORATED HEREIN BY REFERENCE)

Yang G C, Wan L S, Del Priore G. Factors influencing the detection of uterine cancer by suction curettage and endometrial brushing. J Reprod Med 2002; 47:1005-10.

Ries L A G, Melbert D, Krapcho M, Mariotto A, Miller B A, Feuer E J, Clegg L, Horner M J, Howlader N, Eisner M P, Reichman M, Edwards B K (eds). SEER Cancer Statistics Review, 1975-2004, National Cancer Institute. Bethesda, Md., seer.cancer.gov/csr/1975_2004/, based on November 2006 SEER data submission, posted to the SEER web site, 2007.

McCluggage W G. My approach to the interpretation of endometrial biopsies and curettings. J Clin Pathol. 2006; 59:801-12.

accuracy of endometrial sampling in the diagnosis of the patients with endometrial carcinoma and hyperplasia: a meta-analysis. Cancer 2000; 89(8):1765-72.

Feldman S, Berkowitz R S, Tosteson A N. Cost-effectiveness of strategies to evaluate postmenopausal bleeding. Obstet Gynecol 1993; 81(6):968-75.

Grimes D.A. Diagnostic dilation and curettage: A reappraisal. Am J Obstet Gynecol 1982; 142:1-6.

Ong S, Duffy T, Lenehan P, Murphy J. Endometrial pipelle biopsy compared o conventional dilatation and curettage. Jr J Med Sci 1997; 166:47-9.

Tahir M M, Bigrigg M A, Browning J J, Brookes S T, Smith PA. A randomized controlled trial comparing transvaginal ultrasound, outpatient hysteroscopy and endometrial biopsy with inpatient hysteroscopy and curettage. Br J Obstet Gynecol 1999; 106(12):1259-64.

Ferry J, Farnsworth A, Webster M, Wren B. The efficacy of the pipelle endometrial biopsy in detecting endometrial carcinoma. Aust N Z J Obstet Gynecol 1993; 33:1-76.

Guido R S, Kanbour-Shakir A, Rulin M, Christopherson W A. Pipelle endometrial sampling: sensitivity in the detection of endometrial cancer. J Reprod Med 1995; 40:553-5.

Stovall T G, Photopulos G J, Poston W M, Ling F W, Sandles L G. Pipelle endometrial sampling in patients with known endometrial carcinoma. Obstet Gynecol 1991; 77:954-6.

Van den Bosch T, Vandendael A, Wranz P A, Lombard C J. Endopap-versus Pipelle-sampling in the diagnosis of post- 5 menopausal endometrial disease. Eur J Obstet Gynecol Reprod Biol 1996; 64:91-4.

Huang G S, Gebb J S, Einstein M H, et al. Accuracy of preoperative endometrial sampling for the detection of high-grade endometrial tumors. Am J Obstet Gynecol 10 2007; 196:243.e1-243.e5.

Kozuka T. Patch testing to exclude allergic contact dermatitis caused by povidone-iodine. Dermatology 2002; 204 Suppl

Borja J M, Galindo P A, Gomez E, Feo F. Contact dermatitis 15 due to povidone-iodine: allergic or irritant?. J Investig Allergol Clin Immunol 2003; 13(2):131-2.

Naim N M, Mandy Z A, Ahmad S, Razi Z R M. The Vabra aspirator versus the pipelle device for outpatient endometrial sampling. Aust NZJ Obstet Gynecol 2007; 47(2):132-20

Phillips V, McCluggage W G. Results of a questionnaire regarding criteria for adequacy of endometrial biopsies. J Clin Pathol. 2005; 58:417-9.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows the inner obturator and disrupting elements, which may be, for example, a brush, tapered helical screw, loop or loop with brush elements, or the like;

FIG. 2 shows a plunger elements affixed around the obtu-

FIG. 3 shows the obturator within the sheath, with the disruptor retracted and the plunger, which is a ring tightly affixed to the obturator and having close clearance within the 35

FIG. 4 shows three stages of operation of the biopsy device: initially retracted before use, extended to sample the tissue, and retracted with sample drawn into sheath;

FIG. 5 shows a Pipelle endometrial biopsy device of the 40 prior art, which lacks a disrupting device; and

FIG. 6 shows a Tao Brush endometrial biopsy device of the prior art, which lacks an effective suction function; and

FIG. 7 shows a representation of a biopsy procedure of the prior art.

DETAILED DESCRIPTION OF THE PREFERRED **EMBODIMENTS**

A preferred embodiment of the present invention consists 50 of an outer thin walled tube (A) of variable diameter and length. In general dimensions are approximately 1.5 mm outside diameter and 1.4 mm inside diameter; length is between 20-50 cm. This tube may be a clear, malleable plastic tube, such as polyethylene. The inner obturator is preferably 55 formed from a thin wire equivalent of approximately 0.1-0.2 mm diameter, having sufficient mechanical properties to convey the forces for extension and retraction during use. At one end, the one that enters the uterus, a disruptive enhancement is attached to the end. For example, the disruptive enhance- 60 determine infertility causes. ment may have a screw like pattern mechanical attachment, a thin wire or multiple small semi-rigid projections that contact the uterus and loosen and collect the desired specimen. Alternatively, the disruptive enhancement may also include an enzymatic applicator and liquid based digestive enzymes that 65 can be injected through the obturator tube (sheath), for example during extension of the obturator, or through a sec-

10

ond lumen. A tight fitting ring around the obturator acts as a piston and creates the suction as the obturator is withdrawn through the outer tube.

FIG. 1 shows the obturator, which is approximately 0.1-0.2 mm diameter with attached disrupting enhancements of various possible types, and is 45-50 cm long. The obturator is generally stiff, so that the pushing force is transmitted along it's length, within the tube, and thus can be extended and withdrawn. The clearance between the obturator and the inner diameter of the tube is such that a sample volume can be collected within the tube at the distal end, while supporting the obturator to effectively transmit the forces. The tube may have a taper, such that the diameter is greater near the distal end, where the ring moves and disruptive element is extended and retracted, than the proximal end. A plunger handle is provided at the proximal end for manual operation of the extension and retraction.

FIG. 2 shows a tight clearance ring attached to obturator to create suction when withdrawn within the sheath.

FIG. 3 shows the obturator within the sheath, with the disruptor retracted and the plunger, which is a ring tightly affixed to the obturator and having close clearance within the sheath. In this case, the disruptor may be similar to that provided on the Tao Brush $^{TM},$ and, for example, may have a spherical knob at the end to prevent damage to tissue from the tip. Alternately, the disruptor may have bristles at the tip; in any case, a sharp thin tip is generally avoided to limit the risk of puncture.

FIG. 4 shows three stages of operation of the biopsy device. Initially, the obturator is retracted before use, with the disruptive enhancement within the sheath, so that tissue samples are not collected during insertion to the biopsy site. After positioning, the disruptive enhancement is extended from the sheath, to free cellular samples at the biopsy site. Finally, the disruptive enhancement is withdrawn back into the sheath, with the ring creating a suction within the sheath around the obturator to draw a volume of fluid into the sheath. The distal end of the biopsy device may have a greater inner diameter than the proximal end, permitting a larger disruptive enhancement than otherwise possible, and a larger volume of tissue sample to be drawn. The sheath, in that case, may have a constant outer diameter and a tapered wall thickness, a larger outer diameter, or both.

FIG. 5 shows a known Pipelle endometrial biopsy device, 45 which lacks a disrupting device at its distal end. This device has an internal plunger, which creates a suction to draw a sample at the biopsy location.

FIG. 6 shows a known Tao BrushTM endometrial biopsy device, which lacks an effective suction function. Therefore, the amount of fluids surrounding the obturator collected is effectively limited, and the bulk of the sample collected is either attached to the brush at the distal end or is present in the fluids immediately proximal to the brush when it is retracted.

FIG. 7 shows a representation of a biopsy procedure. The detail of the disruptive enhancement according to the present invention is not shown.

The invention may be used, for example, to sample the inside of the uterus to diagnose abnormalities. It can detect or exclude a cancer. It can obtain an adequate tissue sample to

What is claimed is:

- 1. A biopsy device, comprising:
- a flexible coaxial structure, comprising a flexible impermeable sheath and a displaceable flexible tensile and compressive force transmitting structure having a distal end within the sheath, the sheath being configured for

- advancement in direct external contact with tissues into an organ due to a pressure exerted at a proximal end;
- a tissue disruptive sampling structure attached to the displaceable flexible tensile and compressive force transmitting structure at the distal end, configured to preserve three-dimensional structures among cell aggregates of a disrupted tissue; and
- a piston-like structure, disposed around the displaceable flexible tensile and compressive force transmitting structure, disposed proximally with respect to the tissue 10 disruptive sampling structure, configured to slide within the sheath and to draw a vacuum within the sheath and thereby to draw a liquid sample containing the three-dimensional structures among cell aggregates of a disrupted tissue into the sheath from a space surrounding 15 the tissue disruptive sampling structure when the displaceable flexible tensile and compressive force transmitting structure is supplied with a tensile force and the piston-like structure withdrawn into the sheath.
- 2. The biopsy device according to claim 1, wherein the 20 tissue disruptive sampling structure comprises a plurality of bristles.
- 3. The biopsy device according to claim 2, wherein the plurality of bristles extend from the displaceable flexible tensile and compressive force transmitting structure, having an 25 unconstrained diameter greater than a diameter of the sheath, and adapted to be withdrawn into the sheath.
- **4**. The biopsy device according to claim **1**, further comprising a blunt cap at a distal tip of the tissue disruptive sampling structure.
- **5**. The biopsy device according to claim **1**, further comprising an atraumatic bulb at a distal tip of the tissue disruptive sampling structure.
- **6**. The biopsy device according to claim **1**, wherein the displaceable flexible tensile and compressive force transmit- 35 ting structure comprises a guidewire.
- 7. The biopsy device according to claim 6, wherein the guidewire comprises a spirally twisted flexible wire.
- **8**. The biopsy device according to claim **7**, wherein the sampling end comprises a plurality of bristles fixed within the 40 spirally twisted wire at a distal end, which form a tapered brush.
- 9. The biopsy device according to claim 1, wherein the sheath has an outer diameter of between 1 and 3 mm.
- 10. The biopsy device according to claim 1, wherein the 45 sheath comprises at least two lumens, a first lumen containing the displaceable flexible tensile and compressive force transmitting structure, and the second lumen being configured to inject a liquid at a distal end of the sheath.
 - **11**. A biopsy method, comprising: providing:
 - a flexible coaxial structure, comprising a flexible impermeable sheath and a displaceable flexible tensile and compressive force transmitting structure having a distal end within the sheath, the sheath being configured to for 55 advancement in direct external contact with tissues into an organ due to a pressure exerted at a proximal end;
 - a tissue disruptive sampling structure attached to the displaceable flexible tensile and compressive force transmitting structure at the distal end, configured to preserve 60 three-dimensional structures among cell aggregates of a disrupted tissue; and
 - a piston-like structure, disposed around the displaceable flexible tensile and compressive force transmitting structure, disposed proximally with respect to the tissue 65 disruptive sampling structure, configured to slide within the sheath and to draw a vacuum within the sheath and

12

thereby to draw a sample into the sheath from a space surrounding the tissue disruptive sampling structure containing the three-dimensional structures among cell aggregates of a disrupted tissue, when the displaceable flexible tensile and compressive force transmitting structure is supplied with a tensile force and the pistonlike structure withdrawn into the sheath;

inserting the flexible coaxial structure with the tissue disruptive sampling structure contained within the sheath into a body;

- extending the tissue disruptive sampling structure from the sheath and contacting the tissue disruptive sampling structure against a tissue;
- withdrawing the tissue disruptive sampling structure into the sheath, to thereby create the vacuum and obtain the sample; and
- removing the flexible coaxial structure with the tissue disruptive sampling structure contained within the sheath from the body.
- 12. The biopsy method according to claim 11, wherein the sampling end comprises a plurality of bristles which extend from the displaceable flexible tensile and compressive force transmitting structure, having an unconstrained diameter greater than a diameter of the sheath, and adapted to be withdrawn into the sheath.
- 13. The biopsy method according to claim 11, wherein the distal end comprises an atraumatic bulb at a distal tip of the sampling end.
- 14. The biopsy method according to claim 11, wherein the displaceable flexible tensile and compressive force transmitting structure comprises a spirally twisted flexible guidewire, having a plurality of bristles.
- 15. The biopsy method according to claim 11, wherein the sheath comprises a plastic tube having an outer diameter of between 1 and 3 mm.
- 16. The biopsy method according to claim 11, wherein the sheath comprises at least two lumens, a first lumen containing the displaceable flexible tensile and compressive force transmitting structure, and the second lumen being configured to inject a liquid at a distal end of the sheath, further comprising injecting the liquid through the second lumen.
 - 17. A biopsy device, comprising:

50

- a flexible coaxial structure, comprising
- a tubular flexible impermeable sheath configured for advancement in direct external contact with tissues into an organ due to a pressure exerted at a proximal end; and
- a displaceable flexible tensile and compressive force transmitting wire disposed and configured to be displaced axially within the sheath, having at least one state in which an end of the wire extends from an end of the sheath;
- a sampling structure attached to the wire proximate to the end of the wire, configured to disrupt a surface of a tissue, and to preserve three-dimensional structures among cell aggregates of a disrupted tissue; and
- a piston-like structure, attached to the wire proximate to the sampling structure with respect to the end of the wire, configured to draw a liquid sample into the sheath, containing the three-dimensional structures among cell aggregates of a disrupted tissue, when withdrawn into the sheath by a tension on the wire.
- 18. The biopsy device according to claim 17, wherein the sampling structure comprises a plurality of bristles extending outwards from the wire, having an extended diameter greater than a diameter of the sheath, and being adapted to be deflected when withdrawn into the sheath.

19. The biopsy device according to claim 17, further comprising a blunt cap at a distal tip of the sampling end.20. The biopsy device according to claim 17, wherein the sheath comprises at least two lumens, a first lumen containing the wire, and the second lumen being configured for fluid 5

* * * * *